






**Pharmaceutical composition for the oral administration of flavonoids**

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




**Patent number:** EP0711560  
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**Inventor:** HUET DE BAROCHEZ BRUNO (FR); PIOT NOEL (FR);  
CUINE ALAIN (FR)  
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 CY2265 (B)

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 FR2661610  
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 EP0541874

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**Abstract of EP0711560**

Oral compsns. contain a rutaceous flavonoid, partic. micronised diosmin (I), as an effervescent granulate in the form of tablets or sachets. The compsn. pref. comprises 0.1-5 g (I) which is pref. micronised to a particle size of 0.1-10  $\mu$ m. The effervescent formulation is achieved by use of a carboxylic acid and one or more carbonates. Other ingredients include sweeteners, flavourings and carriers. The small amt. of water present in (I) is used to start a single phase granulation process.

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